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ZIMAKOV, A. M. and G. S. BOBROVSKII

Parovye kotly promyshlennykh predpriiatii. Moskva, Gosenergoizdat, 1949. 335 p. diagra.

Steam boilers in industrial enterprises.

DLC: TJ289.B57

SO: Manufacturing and Mechanical Engineering in the Soviet Union, Library of Congress, 1953

ZIMBROY, A.M.

AUTHORS: Vorobey, V.V. and Zimakov, A.M.

133-11-13/19

TITIE:

Automation of Thermal and Technological Processes in Iron and Steel Works (Avtomatizatsiya teplovykh i tekhnologicheskikh protsessov na predpriyatiyakh chernoy metallurgii)

PERIODICAL: Stal', 1957, No.11, pp. 1024 - 1027 (USSR).

ABSTRACT: Automatic controlling used in blast furnaces, openhearth and electric furnaces and rolling mills is outlined. It
is pointed out that the progress achieved is to a large extent
due to the existence of special organisations grouped in the
"Energochermet" Trust: instrument-making factories ("KIP" in
Kharkov and "Teplopribor" in Chelyabinsk), the Central Laboratory
of Automatics. Central Design Office. etc.

AVAILABLE:

Library of Congress

Card 1/1

"Stream Boilers in Industrial Enterprises". Gosenergoi dat, Moscow/ Leningrad, 1969, 331 pp, 11 rubles h0 kopeks.

SO: W-1h151 11 Oct. 1950.

ZIMAKOV, B.M.; STEPANOV, Yu.V.

Oil manifestation in the Vorkuta coal-bearing region of the Pechora Basin. Sov. geol. 8 no.3:125-127 '65.

(MIRA 18:5)

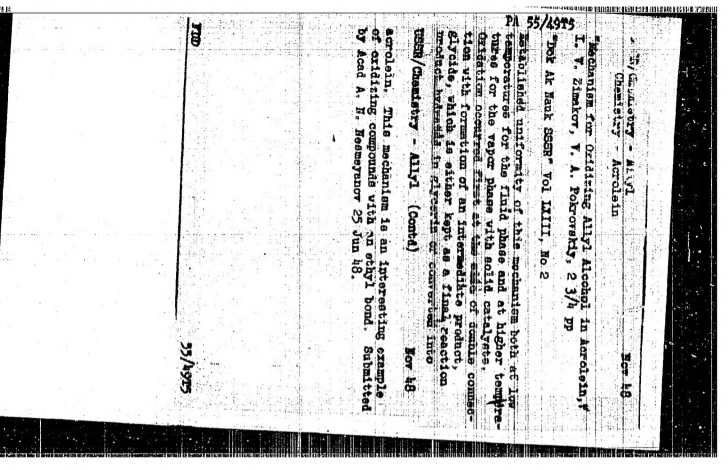
1. Vorkutskaya kompleksnaya geologorazvedochnaya ekipeditsiya UTGU i Moskovskiy geologorazvedochnyy institut im. S. Ordzhonikidze.

ETTINGER, I. L.; YEREMIN, I. V.; ZIMAKOV, B. M.; BAKALDINA, A. P.

Sorption properties of various petrographic components of fossil coals. Dokl. AN SSSR 155 no. 2:364-367 Mr '64. (MIRA 17:5)

1. Institut gornogo dela im. A. A. Skochinskogo, Moskovskiy geologorazvedochnyy institut im. S. Ordshonikidze i Institut geologii i razrabotki goryuchikh iskopayemykh. Predstavleno akademikom N. V. Mel nikovym.

(Pechora Basin—Mine gases)		Effect of geoldistribution of Izv.vys.ucheb.: 1. Moskovskiy Ordzhonikidze.	av.;gec	in coal depo l.i razv. 4	sits of no.7:78-	the Peci	hora Ba	asin. (14:8)	
			(Pechor	a Basin-Min	e gases)				
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SPITSYN, VIKT, I., akademik; ZEMLYANOVA, L.I.; MIKHAYLENMO, I.Ye.;

Electron microscope study of the effect of radioactive waves from solids on the structure of their surface. Dokl. AN SSSR 139 no.5:1163-1165 Ag. '61. (NIRA 14:8)

1. Institut fizicheskoy khimii AN SSSR. (Solids, Effect of radiation of)

ZIMAKOV, I.Ye.; DORROSEL'SKAYA, N.P.; SPITSYN, Vikt.I., akademik

Effect of the radioactivity of molybdic andydride on the variation of its specific surface area. Dokl.AN SSSR 148 no.4:884-885 F *63. (HIRA 16:4)

1. Institut fizicheskoy khimii AN SSSR. (Nolybdenum oxides) (Radioactivity) (Sutface measurement)

ACCESSION NR: AP4020063

S/0186/64/006/001/0130/0132

AUTHOR: Spitsy*n, Vikt. I.; Zimakov, I. Ye.; Dobrosel'skaya, N. P.

TITLE: Effect of radioactive emission S35 and Mo⁹⁹ on the magnitude of the specific surface of molybdenum disulfide

SOURCE: Radiokhimiya, v. 6, no. 1, 1964, 130-132

TOPIC TAGS: molybdenum disulfide, specific surface, BET method, radioscitive radiation, tagged molybdenum disulfide, crystal lattice irregularity, crystallization center formation, specific surface change, sulfur 35, molybdenum 99

ABSTRACT: The effect of rad cactive radiation imparted by incorporating \$35 and Mo⁹⁹ isotopes, on the specific surface of MoS₂ obtained from a melt was investigated. The specific surface was determined by the BET method by low temperature adsorption of kryston. The specific surface of the radioactive materials differs from that of the non-tagged MoS₂ in that it increases with an

Card 1/3 .

ACCESSION NR: AP4020063

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increase in specific radioactivity. With \$35 the increase is fairly regular and the specific surface is tripled with 30 millicurie/gm. radioactivity from about 8 to 24 m²/gm. With Mo³9 the increase is sharper, i.e., almost tripled with 15 millicurie/gm. radiation. fig. 1). The specific surface of the materials does not change on standing. The difference in specific surface associated with radioactive radiation is apparently caused by the formation of irregularities in the crystal lattice and the creation of additional crystallization centers. Orig. art. has: 2 figures and 1 table

ASSOCIATION: None

SUBMITTED: 05Aug63

SUB CODE: GP, NP

DATE ACQ: 31Mar64

r64 ENCL: 01

NO REF SOV: 007

OTHER: 003

Card 2/3

ZIHAKOV, I. Ye.

The Second All-Union Conference on the Preparation and Analysis of High-Purity Elements, held on \$4-28 December 1963 at Gorky State University im. N. I. Lobachevskiy, was sponsored by the Institute of Chemistry of the Gorky State University, the Physicochemical and Technological Department for Inorganic Materials of the Academy of Sciences USSR, and the Gorky Section of the All-Union Chemical Society im. D. I. Mendeleyev. The opening address was made by Academician N. M. Zhavoronkov. Some 90 papers were presented, among them the following:

A. A. Popel' and Z. A. Saprykovo. Quantitative determination of paramagnetic ions in solution by NMR methods.

I. Ye. Zimakov. Determination of microimpurities (10⁻⁷ to 10⁻¹⁸%) by repeated radioactive dilution.

A. A. Tumanov and V. S. Yefimychev. Determination of micro-concentrations with salicylan-2-aminophenol.

(Zhur ANAL Khim, 19 NO.6, 1964, p.777-79)

8/137/62/000/012/082/085 A006/A101

AUTHOR:

Zimakov, I. Ye.

TITLE:

The use of radioactive tracers to check some analytical methods of separating rhenium and molybdenum

PERIODICAL:

Referativnyy zhurnal, Metallurgiya, no. 12, 1962, 15, abstract 12K92 ("Sb. nauchn. tr. Oos. n.-i. in-t tsvetn. met.", 1961, no. 18, 56 - 59)

TEXT: Using radioactive Mo⁹⁹ and Re¹⁸⁶ isotopes, the author compares 2 methods of separating low Re amounts from high Mo amounts with the aid of 8-axyquinoline and x-benzoin oxime. The oxyquinoline method was found to be suitable for Re separation from Mo even if Mo : Re = 585,000 : 1. Most complete Mo separation from Re by chloroform extraction is obtained at pH 4.5. Using Re¹⁸⁰, it was established that during extraction Re is not transferred completely into the chloroform layer. The α-benzoin oxime method can not be used to separate low Re from high Mo amounts, due to considerable sorption of Re by a precipitate of Mo α-benzoin oxime. There are 6 references.

[Abstracter's note: Complete translation]

L. Vorob'yeva

Card 1/1

ZEMLYANOVA, L.I.; ZIMAKOV, I.Ye.; LYAPINA, A.M.; SPITSYN, Vikt. I.;

Electron microscope study of the effect of radioactivity of elementary sulfur on the structure of its surface.

(Radiokhimiia 5 no.3: 192-394 '63. (MIRA 16:10)

(Sulfur isotopes) (Electron microscopy)

SOV/136-59-5-15/21

AUTHORS: Gulyayeva, Ye.I., Zimakov, I.Ye., and Rudenko, B.I.

TITLE: Extraction of Rhenium from Industrial Solutions using

Activated Coal (Izvlecheniye reniya iz proizvodstvennykh

rastvorov pri pomoshchi aktivirovannykh ugley)

PERIODICAL: Tsvetnyye metally, 1959, Nr 5, pp 73-77 (USSR)

ABSTRACT: The difficulty in extracting Re from solution is the separation from W and Mo which have similar properties.

An ordinary chromatographic method was tried using activated coals types KAD, SK-T, SU-KhU, and MSK-1. The coals were washed with 0.01N H2SOL until they showed

acid reaction to methyl orange and then the test solutions were poured through the coal. Afterwards the

coal was washed with water and then 1% soda solution to remove the Re, W and Mo. The solutions used are given in Table 1. Radicactive isotopes Reloo, Mo99, and W185

were added to the solutions and used to indicate the degree of separation. The best coal was found by testing

with the first solution. 0.3, 0.5 and 1.0 g of coal and 4.5 and 7.0 mm diameter columns were tried. Table 2

Card 1/3 shows the dynamic exchange capacity under various conditions. It increases with increase in the ratio

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Extraction of Rhenium from Industrial Solutions using Activated Coal

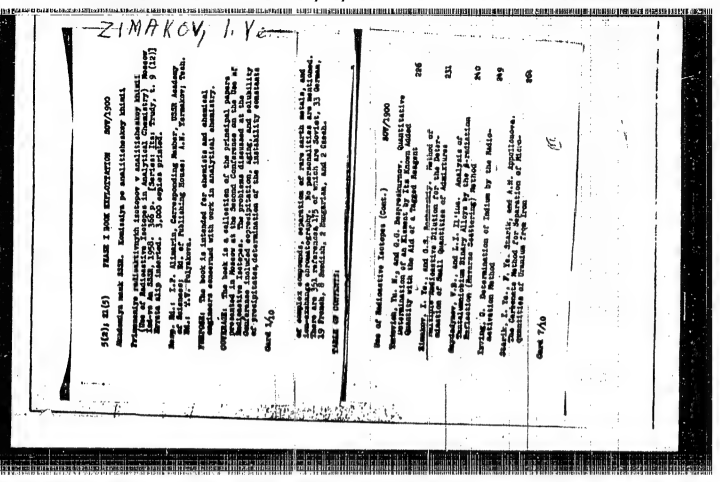
height/diameter of column. Fig 2 shows that MSK coal has the greatest absorption. The washing results in Fig 3 show that Re is completely extracted from all the coals so MSK coal was used for further work with the other three solutions. Table 3 shows that the exchange capacity was highest for solution 2 which differed from the other solutions in Mo content and acidity. shows that Re is selectively absorbed by the coal but Mo and W are weakly absorbed. Tests on acidity showed that absorption of W and Mo were practically independent of pH value, but Re absorption increases with inverse pH. Therefore washing was carried out with alkaline solution. Fig 8 shows that Mo and W are washed out before Re. The final solution containing Re has small amounts Mo and

Card 2/3 W present. Hot soda gave better results than cold soda.

Extraction of Rhenium from Industrial Solutions using Activated Coal Further research to obtain more concentrated and purer Re is being carried out.

There are 8 figures, 4 tables and 1 Soviet reference.

Card 3/3



ZIMAKOV, I.Ye.; SPITSYN, Vikt.I., akademik

Effect of radioactive radiation energy on the rate of evaporation of a solid. Dokl. AN SSSR 141 no.6:1400-1402 D '61. (MIRA 14:12)

1. Institut fizicheskoy khimii AN SSSR.
(Radioactivity) (Evaporation)

SPITSYN, Vikt.I., akademik; ZIMAKOV, I.Ye.

Effect of the radicactivity of molybdenum trioxide on the rate of its vaporization. Dokl. AN SSSR 139 no.3:654-657 Jl *61. (MIRA 14:7)

1. Institut fizicheskoy khimii AN SSSR.

(Molybdenum oxide) (Solids, Effect of radiatom on)

(Vaporization)

S/020/61/141/006/018/021 B105/B147

5 Li 600 AUTHORS:

Zimakov, I. Va., and Spitsyn, Vikt. I., Academician

TITLE:

Effect of radioactive energy on the evaporation rate of a

solid

PERIODICAL: Akademiya nauk SSSR. Doklady, v. 141, no. 6, 1961, 1400-1402

TEXT: This is a supplement to a previous paper by the authors on the effect of the radioactive level upon the evaporation rate of MoO, (Ref. 1: Vikt. I. Spitsyn, I. Ye. Zimakov, DAN, 139, No. 3 (1961)), where also the production methods of MoO, with different Mo9 contents were described. The effect of β -radiator additions such as a) γ^{90} (E = 2.18 MeV), b) Mo9 (E = 1.23 MeV), and c γ^{185} (E = 0.43 MeV) as well as the value of their radiant energy were studied. In case a), MoO, powder was soaked with a certain quantity of dissolved radioactive yttrium nitrate (or of not radioactive yttrium nitrate as control), dried, and therefrom Card 1/3

Effect of radioactive energy. ...

S/020/61/141/006/018/021 B103/B147

fractions of 0.5 - 0.25 mm were obtained by screening. The specific radioactivity was related to MoO₃. c) As admixture, radioactive ammonium tungstate was added to the ammonium molybdate solution. After evaporation

tungstate was added to the ammonium molybdate solution. After evaporation of the solution, the residue was moistened with some drops of ${\rm HFO}_3$.

calcined, and fractions obtained like in case a). The evaporation rate was determined in an electric oven from the change in the length of the quartz spring to which the specimen was fixed. A stream of dry air or nitrogen (flow rate 10 ml/min) was passed through at 700 ± 1°C. The changes in the weight of the weighed portions were noted and used for plotting evaporation curves. The evaporation rate (ER) of the preparations containing non-radioactive Y and of those with an initial specific radioactivity of 1.0 and 2.0 millicuries/g is practically constant. From 3 millicuries/g onward, the ER of the non-radioactive specimens differs considerably. The ER of each specimen increases steadily, since the radioactivity of the residual preparation increases owing to MoO₃

evaporation. In the case of a) this difference sets in at 5 milliouries/g. No radioactive products were found in the sublimate. The dependence of the ER of MoO, on its specific radioactivity does not differ from that Card 2/3

Effect of radioactive energy...

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described in Ref. 1. Thus, it has been found that the ER of MoO3 depends not only on the radioactivity level of the preparation, but also on the radiant energy of the radioactive addition. The ER of MoO, is affected not only by the radiation of a) but also by the radiation of the foreign radioactive inclusions in the solid phase. There are 4 figures and 1

ASSOCIATION: Institut fizicheskoy khimii Akademii nauk SSSR (Institute of Physical Chemistry of the Academy of Sciences USSR)

SUBMITTED:

July 22, 1961

Card 3/3

ZIMAKOV, I.Ye.; ROZHAVSKIY, G.S.

Method of multiple radioactive dilution for the determination of small quantities of admixtures. Trudy kom.anal.khim. 9:231-239 [58. (NIRA 11:11)

(Radiochemistry)

5(2) AUTHOR:

Zimakov, I. Ye.

SOV/32-25-2-2/78

TITLE:

The Determination of the Solubility of Barium-tungstate by Means of the Radioactive Tungsten Isotope - 185 (Opredeleniye rastvorimosti vol'framata bariya s pomoshch'yu radioaktiv-nogo izotopa vol'frama - 185)

PERIODICAL:

Zavodskaya Laboratoriya, 1959, Vol 25, Nr 2, pp 133-134 (USSR)

ABSTRACT:

Slightly soluble barium tungstate (I) is used in separating tungsten from rhonium in the determination of the rhenium content in tungsten concentrates. In the present case the solubility of (I) was determined in water. The indicator used (W¹⁸⁵) possesses a \$\beta\$-radiation with an energy of 0.430 MeV and a half-life of 73.2 days. Tests were carried out in a

and a half-life of 73.2 days. Tests were carried out in a vessel with a spiral agitator. The activity of the dry residue was measured by means of the counter MST-17 and the solubility calculated on the basis of an equation (Table). The curve representing the solubility as a function of the temperature (Fig 1) shows that (I) is the least soluble at 10° C and that the solubility increases moderately at temperatures below 10° C. The dependence of the solubility

Card 1/2

The solubility

The Determination of the Solubility of Bariumtungstate by Means of the Radioactive Tungsten Isotope - 185

on the pH value was studied with a pH meter of the LP-5 type with a glass electrode at 20° in hydrochloric acid and soda lye. The investigation results show (Fig 2) that the precipitation of (I) is most complete at a pH value of 6-7. There are 2 figures and 1 table.

ASSOCIATION:

Gosudarstvennyy nauchno-issledovatel'skiy institut tsvetnykh metallov (State Scientific Research Institute for Non-ferrous Metals)

Card 2/2

SPITSYN, Vikt.I., akademik; ZIMAKOV, I.Ye.

Effect of radioactive radiation from Y90 and Y91 on the solubility of yttrium hydroxide. Dokl.AN SSSR 138 no.1:130-132 My-Je '61. (MIRA 14:4)

1. Institut fizicheskoy khimii AN SSSR.
(Yttrium--Isotopes) (Yttrium hydroxide) (Solubility)

Use of radioactive tracers to check certain analytical methods of separating rhenium and molybdenum. Scor. nauch. trud. Gintsvetmeta no.18:56-59 '61. (MIRA 16:7) (Rhenium—Metallurgy) (Molybdenum—Metallurgy) (Radioactive tracers)

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\$/020/61/139/003/024/025 B127/B206

AUTHORS:

Spitsyn, Vikt. I., Academician, and Zimakov, I. Ye.

TITLE:

Effect of radioactivity of molybdenum anhydride on the rate

of its evaporation

PERIODICAL:

Akademiya nauk SSSR. Doklady, v. 139, no. 3, 1961, 654-657

TEXT: For their studies the authors used preparations of molybdenum anhydride containing Mo 99 (half-life period 68.3 hr) which emits β -particles of high energy (E $_{max}$ %1.23 MeV, 85 %). The specimens had a specific activity of 1.0-28.0 millicuries/g. A mixture of calculated amounts of highly active and inactive MoO3 was converted with ammonia into ammonium molybdate; the solution was evaporated and annealed with small amounts of nitric acid to prevent the reduction of hexavalent Mo through ammonia. The MoO3 obtained was mixed with small amounts of Nb205 and ZrO2, and sublimated at B50°C in a weak air current for purification from radiochemical admixtures. The particle size was 0.25-0.5 mm. Card 1/5

Effect of radioactivity of molybdenum...

S/020/61/139/003/024/025 B127/B206

Working method: A dry current of air was conducted through the layer of molybdenum powder at a rate of 8 ml/min and 700°C temperature. Every 15 min, the quartz crucible was removed, cooled, weighed, and again suspended in the tube. Table 1 shows the experimental results. The following dependence between rate of evaporation and radicactivity was established: At a specific activity of MõO₃ of 2.5-3 millicuries/g, the

rate of evaporation equals the inactive specimen. At a specific activity of 2.5-4 millicuries/g, a drop of the rate of evaporation was established. At 4-9 millicuries/g, a fast increase of the rate of evaporation was observed, which was retarded at a further increase of the specific activity. An increase of the electric charge on the surface was observed on the basis of β -particle emission. It probably also had an effect on the rate of evaporation, which becomes obvious in an interaction between the surface of the hard substances and the gaseous molecules. In the MoO₂ molecule, the Mo is surrounded by three negatively charged oxygen atoms. The positive charge of the solid MOO₂ prevents, therefore,

transition into the vapor phase. The following experiment was made for explanation: In a crucible with radioactive MoO3, an earthed platinum

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Effect of radioactivity of molybdenum...

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wire was brought directly to the specimen. The results in Figs. 3 and 4 confirm the assumption that the electric charge causes the retardation of the rate of evaporation at 2.5-4 millicuries/g. The increase of the rate of evaporation of MöO, above 4-5 millicuries/g depends on the radiation effects in the gaseous phase. A strong flow of β-particles ionizes the evaporated molecules to MoO, which repel themselves from the surface. The surface of the crystals is changed through radiometric radiation. The electron-microscopic pictures showed a surface for MoO, entirely different from that for MŏO, which is full of irregularities, elevations, and depressions, the size of which depends on the strength of the specific radioactivity. It is also pointed out that the rate of evaporation begins to increase only at dose > 10.16 eV/g·sec in the case of an external irradiation of MoO, by an electron current of 800 kev. There are

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Effect of r	adioactivity	of moly	bdenum	3	3/020/0 1127/B2	61/139) 206	/003/024/04	25	V
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SUBMITTED:	April 12,	1961		• •		••			
Table 1. Ratevaporation	of MoOz				184				
as a function specific rad of preparati	lioactivity ina	Удельи, радновк- тиви., иСи/г	Колич. непарнями, за 2 часа МоО, (среди,), г	ЧНаблю- давш. от- илон! от среди, %	A mpena- para	Удельи. Видковк- тиви., МСЦ/г	3 Колич. испяриви 84 2 чясь МоО, (среди.), г	HAGAR ABBIN. C MAGN. (CPEAN.	
Legend: (1) of the prepar (2) specific activity. mi	Number cation; tradio-	Неак- тивный 1,0 2,0	0,1240 0,1240 0,1240	5,0 6,0 5,1	7 8 9 10	5.0 7.0 8,4	0,1148 0,1458 0,1514	3,5 5,0 7,0	
(3) amount of value) evapor 2 hr, grams; deviation fr	MoO ₃ (mean 5) rated in 600 (4) observed	2,5 3,0 4,0	0,1217 0,1184 0,1125	5,0 5,0 5,0	10 41 12 .13	10,0 14,0 20,0 28,0	0,1556 0,1600 0,1623 0,1009	4,5 5,0 6,0 6,0	
mean value, Card 4/5	%.	•.							
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S/020/61/139/005/020/021 B103/B208

5.4600 AUTHORS:

Spitsyn, Vikt. I., Academician, Zemlyanova, L. I.,

Mikhaylenko, I. Ye., Gromov, V. V., and Zimakov, I. Ye.

TITLE:

Electron-microscopic examination of the effect of radioactive radiation of solids on the structure of their surface

PERIODICAL: Akademiya nauk SSSR. Doklady, v. 139, no. 5, 1961, 1163-1165

TEXT: The crystal lattice of solids is disturbed by the ionizing action of their own radioactive radiation and the appearing record atoms, which also changes their surface structure. According to the authors, all this may be one of the causes of the effect exerted on physicochemical properties of solids by their own radiation (sorptive power, solubility in water, kinetics of heterogeneous processes of isotopic exchange, catalysis, etc.). The authors made electron-microscopic studies of the surface structure of radioactive samples of K2SO4, MgSO4, BaSO4 and MoO3 which had been used previously to study adsorption, catalysis, and isotopic exchange. Except for BaSO4, the pictures were obtained by Card 1/5

Electron-microscopic examination ...

S/020/61/139/005/020/021 B103/B208

replication, and for BaSO, the method of double replicas (silver-quartz replicas) was used. K_2SO_4 , M_8SO_4 , and MoO_5 were applied to a collection film in the form of a fine powder. A 200 - 300 Å thick quartz layer was sputtered onto it in vacuo. After dissolution of collodion in amylacetate, the quartz replica were rinsed in distilled water in the case of K2SO4 and MgSO4, and in dilute alcohol in the case of MoO4. Radicactive samples of K and Mg were obtained by adding small amounts of NagSO4 containing S35. BaSO4 precipitates were isolated by a method previously described by Vikt. I. Spitsyn, V. V. Gromov (DAN, 123, 722 (1958); Radiokhimiya, 1, 181 (1959)). Radioactive Moo, was obtained by adding an Mo99 containing sample to ordinary Mo03 in order to attain the necessary specific radioactivity. The mixture was converted to ammonium molyboate by treating it with aqueous ammonia; it decomposed when heated. The resultant Moo, was sublimed at 850°C. When comparing the pictures (magnification: 12,000 times) [Abstracter's note: Not reproducible] Card 2/ 5

Electron-microscopic examination ...

S/020/61/139/005/020/02; B103/B208

authors found the following differences in the crystal surface of a radioactive and b) non-radioactive samples: 1) The surface of b) is comparatively smooth, that of a) highly pitted. The crystal surface of BaSO, is changed to a high extent by incorporation of small radium $\mathrm{K}_{2}\mathrm{SO}_{4}$, BaSO_{4} , and MoO_{3} also show some changes in their surface amounts. structure after an external arradiation with 800-key electrons. Although the dose was much higher in this case, the changes were less pronounced. than those caused by radioactive radiation. The above surface defects appear rather regularly over the whole length of the crystal of the radioactive substance. The deep cavities observed in samples irradiated with neutrons were absent. The surface changes resemble those observed in metal etched by an ion beam. The authors further conclude from the comparison of the photographs that the surface defects of the radicactive samples develop already during the separation of the solid phase from the solution or from the gas. They assume that the radiation of electrons or other charged particles during the crystallization of solid substances gives rise to a great number of new active centers (seed crystals). The particle-size distribution on separation of radioactive salts from

Card 3/5

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Electron-microscopic examination...

\$/020/61/139/005/020/021 B103/B208

ASSOCIATION: Institut fizicheskoy khimii Akademii nauk SSSR (Institute of Physical Chemistry of the Academy of Sciences USSR)

SUBMITTED: April 15, 1961

Card 5/5

ZIMAKOV, I. Ye., and ROZHAVSKIY, G. S.

"Method of repeated radicactive dilution for the determination of small admixtures (of the order of 10" to 10" 8)."

report presented at The Use of Radioactive Isotopes in Analytical Chemistry, Conference in Moscow, 2-4 Dec 1957

Yestnik Ak Nauk SSSR, 1958, No. 2, (author Rodin, S. S.)

Determining the solubility of barium tungstate with radicactive tungsten185. Zav.lab. 25 no.2:133-134 '59. (MIRA 12:3)

1. Gosudarstvennyy nauchno-issledovatel'skiy institut tavetnykh
metallov. (Barium tungstates) (Solubility) (Tunguten-Isotopes)

AUTHORS:

Zimakov, I. Ye., Rozhavskiy, G. S.

SOV/32-24-8-2/43

TITLE:

The Method of Multiple Radioactive Dilution for Determining: Trace Amounts in Mixtures (Metod mnogokratnogo radioaktivnogo razbavleniya dlya opredeleniya malykh primesey)

PERIODICAL:

Zavodskaya Laboratoriya, 1958, Vol. 21, ir 3, pp. 922-925 (USSR)

ABSTRACT:

Previous methods of using radioactive dilution for determining trace amounts in mixtures have had a certain disadvantage. The specific activity and the amount of the isolated substance have had to be determined by an analytical method. In this paper a method is described which climinates this disadvantage. This determination involves adding two different amounts m₁ and m₂ of the radioactive isotope of the substance x being determined to two similar solutions of this substance. By withdrawing equal amounts of substance y the concentration of the substance x to be analysed can be calculated according to the activity. The paper gives the formula for calculating the concentration, and gives several other equations as well. If the weighed amount of the sample to be analysed and the apecific activity of the preparation are increased the ad-

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The Method of Multiple Radioactive Dilution for Determining Trace Amounts in Mixtures

SOV/32-24-8-2/43

mixture can be determined in the order of magnitude of 10^{-5} to 10^{-7} 5 with an accuracy of about 10 5. A method of determining trace amounts of antimony in lead was worked out. The main problem here was to find a way by which small and equal amounts of the substance to be analysed could be separated from solutions of different concentrations. The most suitable method found for this purpose was extraction of the autimony-methylviclet complex compound with toluene. The optimal acidity of the antimony solution was found to be 1:9 in HCl. A procedure as well as tables of results obtained are given.

There are 2 tables.

ASSOCIATION

Gosudarstvensky nauchno-issledovatel'skiy institut banetnykh metalky State Scientific Research institute for Non-ferrous Metals)

Card 2/2

S/020/63/148/004/022/025 B144/B101

AUTHORS:

Zimakov, I. Ye., Dobrosel'skaya, N. P., Spitsyn, Vikt. I.,

Academician

TITLE:

Effect of the radioactivity of molybdenum trioxide on the

change of its specific surface

PERIODICAL:

Akademiya nauk SSSR. Doklady, v. 148, no. 4, 1963,

884-885

TEXT: The specific surface was studied in MoO₃ samples containing different quantities of Mo⁹⁹ ($T_{1/2} = 68.3$ hr, E (β) max = 1.23 MeV). A mixture of radioactive and non-radioactive MoO₃ was converted by NH₄OH to ammonium molybdate. The solution was evaporated and the residue calcined with addition of HNO₃ to prevent reduction of sexivalent Mo by NH₃. The MoO₃ recovered was mixed with small quantities of Nb₂O₅ and ZrO₂ and sublimated at 850°C. The surfaces were measured in samples of

Card 1/2

Effect of the radioactivity of

0.5 - 0.25 mm granulation.

S/020/63/148/004/022/025

The external specific surface (without blind pores) was measured by filtering rarefied air through the sample and determining the resistance. At radioactivities up to \$\approx 10 mcu/g, these surfaces decreased slightly, whereas with higher mcu values they increased. The total surface was determined by the BET method with adsorption of krypton (area occupied by the Kr molecule: 19.5 A). Additions of radioactive MoO, up to 10 mou reduced the total specific surface (maximum 30%), while higher additions increased it. The absolute values obtained by the two methods were rather similar, which indicates only a small number of blind pores. Based on a previous study (DAN, 139, 654 (1961)), the reduction of the specific surface with low radioactivities is attributed to the reduction of the evaporation rate. Higher radioactivity entails higher evaporation rates and formation of dendrites owing

to the effect of β -particles. There are 2 figures. ASSOCIATION:

Institut fizicheskoy khimii Akademii nauk SSSR (Instituto

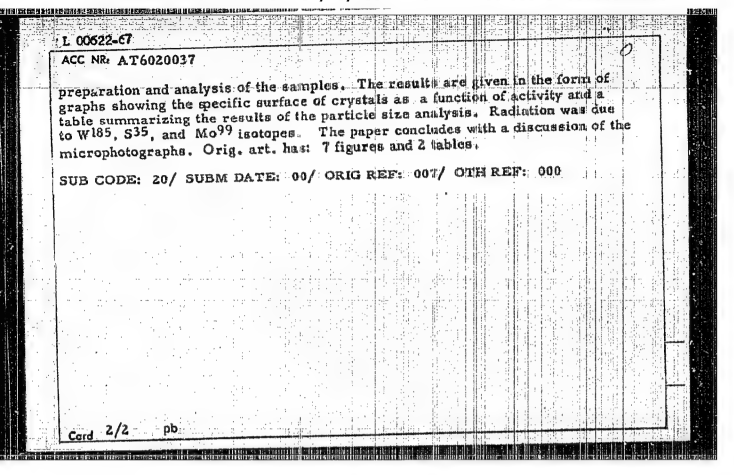
of Physical Chemistry of the Academy of Sciences USSR)

SUBMITTED:

November 16, 1962

Card 2/2

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- 14	ACC NR. AT6020037 (A) SOURCE CODE: UR/2564/65/005/000/0212/0218	
	AUTHOR: Spitsyn, V. I.; Zimakov, I. Ye.; Dobrosel'skaya, N. P. 52	
	ORG: none	
	TITLE: The influence of radiation on the formation and uniformity of crystals	
	SOURCE: AN SSSR. Institut kristallografii. Rost kristallov, v. 5, 1965, 212-218	
	TOPIC TAGS: crystal growth, crystal deformation, radiation damage, radiation	
-	ABSTRACT: In the last 5 years the institute of Physical Chemistry, Academy of Sciences SSSR (Institut fizicheskoy khimii Akademii nauk SSSR) gathered numerous	
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	present article contains further data collected during the state of mains. The	9
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i	precipitation from alloys. The nonradioactive and radioactive samples with differ-	
	were studied by measuring their size and their specific specific and their specific	_
	electron-miscroscopic pictures. The paper describes in considerable detail the	_
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ACC NR: AFFOLIA74

SOURCE CODE: UN/OUTO/66/011/002/0516/0120

AUTHOR: Spitsyn, V. I.; Zimakov, I. Ye.; Zenlyanova, It I.

30

ORG: Institute of Physical Chemistry, Academy of Sciences, MISR (Institut fizicheskoy khimii Akademii nauk SSSR)

TITLE: Investigation of the influence of radioactive calds on the furface structure of molybdamam analydride

SOURCE: Kristallografiya, 2711, mo. 2, 1966, 516-320

TOPIC TAGS: molybdenum compound, radioactivity effect, surface property, civital-

ABSTRACT: The authors investigated under an electron microscope the surface structure of samples of molybdenum tricuide containing different amounts of radioactive Mo^{99} , which emits high-energy β particles (1.23 MeV) of relatively short half life

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ZIMAKOV. N. V.

"Prospects of Using Fission Product Source Radiation in Radiation Chemistry", by N. V. Zimakov, E. V. Volkova, A. V. Fokin, V. V. Kulichenko, V. G. Vereskunov, A. G. Bykov, and N. I. Bogdanov

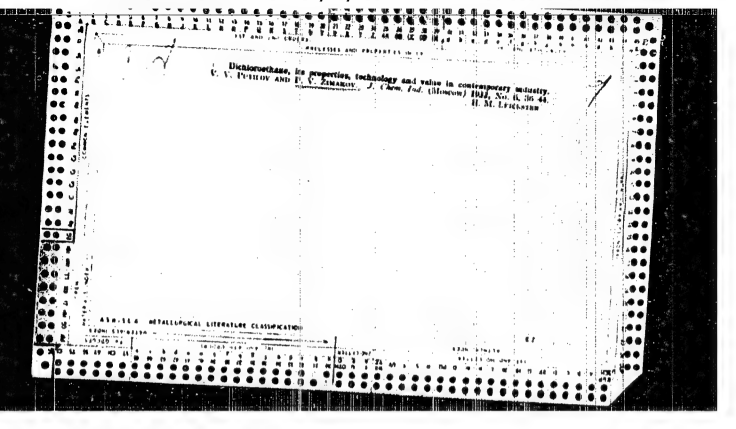
Report presented at 2nd UN Atoms-for-Peace Conference, Geneva, 9-13 Sept 1958

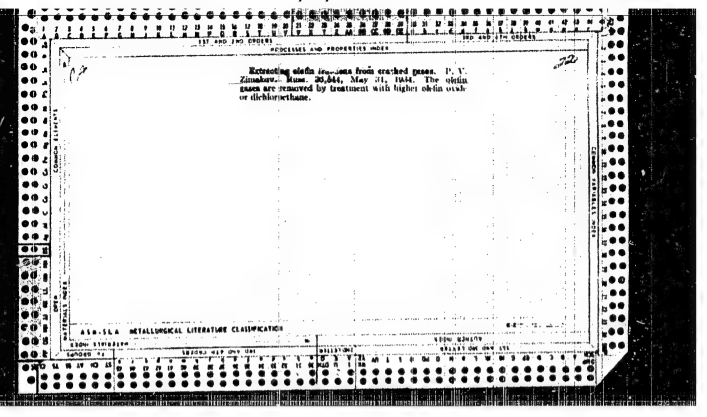
ZIMAKOV, P. P. (Assist. Prof.)

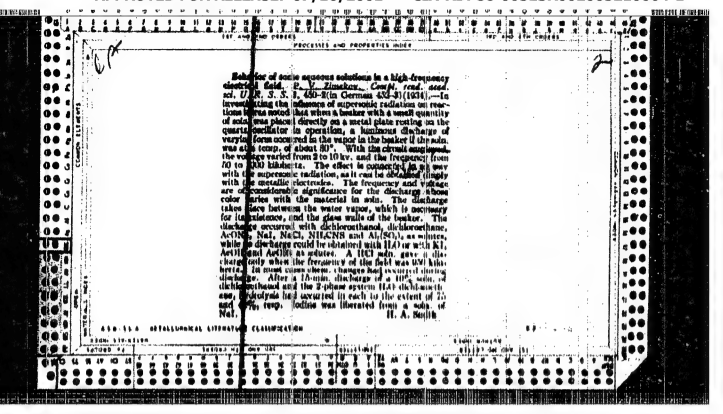
"Trainer for Fighter Pilots."

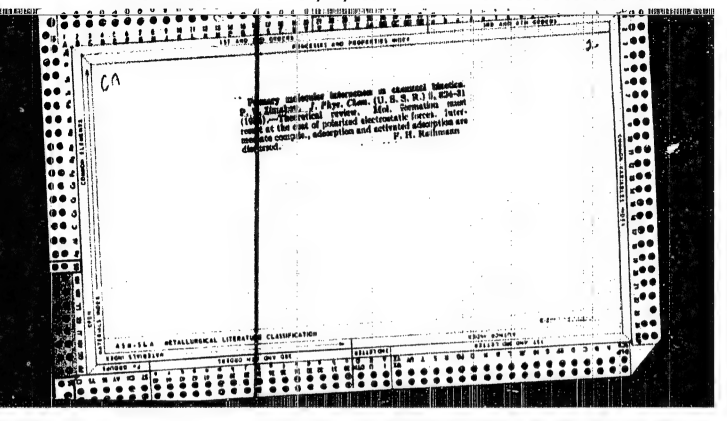
report presented at the 13th Scientific Technical Conference of the Kuybyshov Aviation Institute, March 1959.

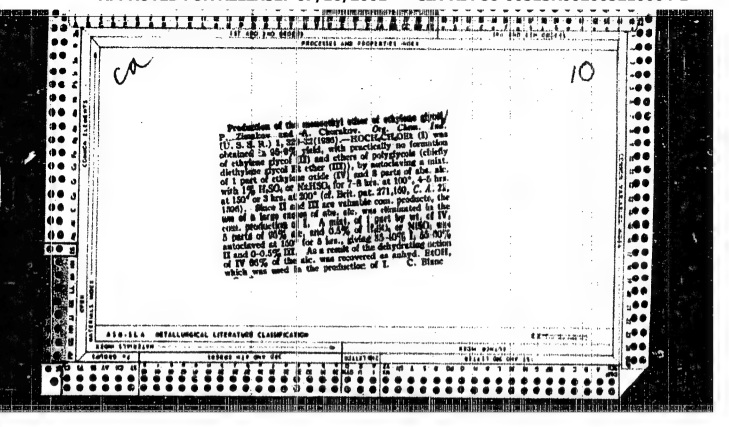


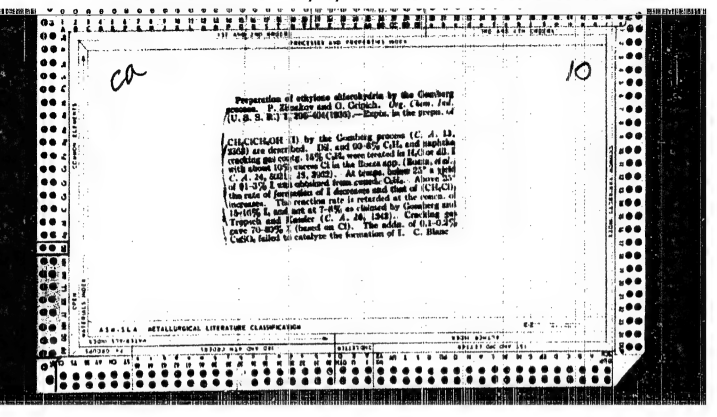


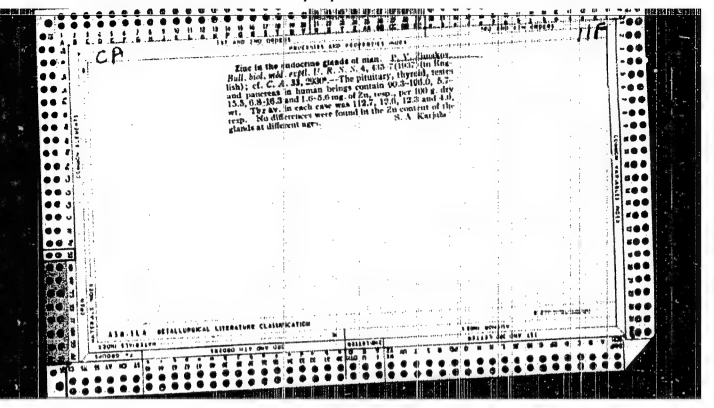


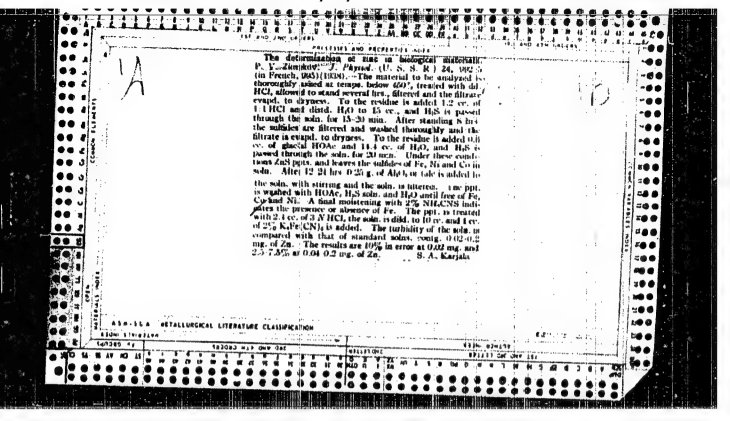


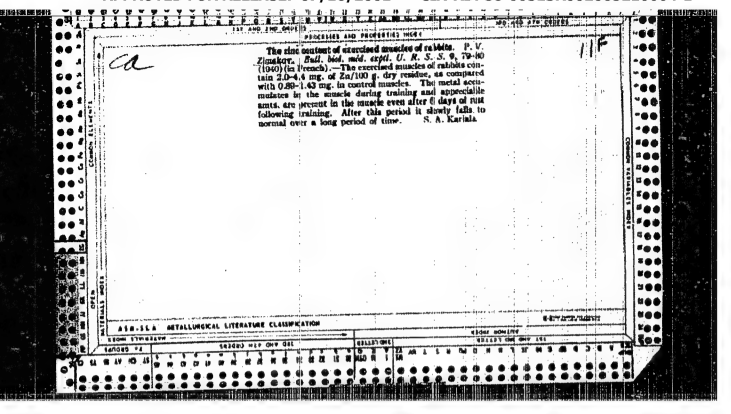


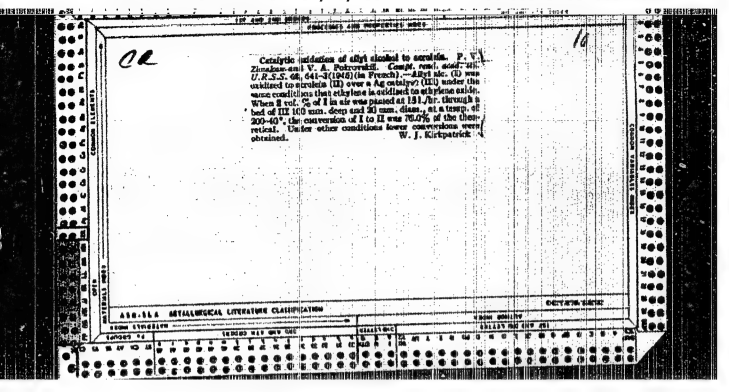


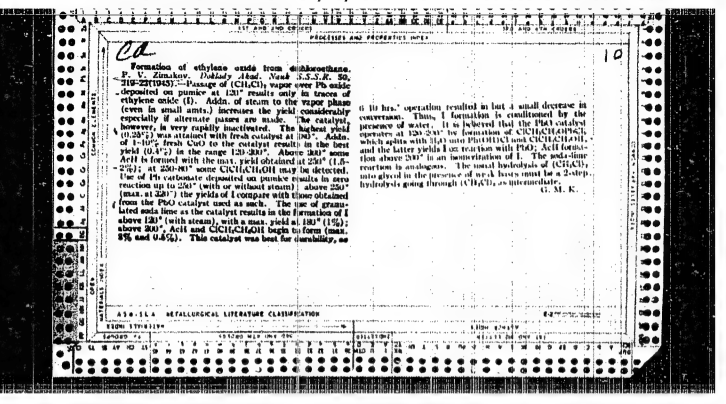


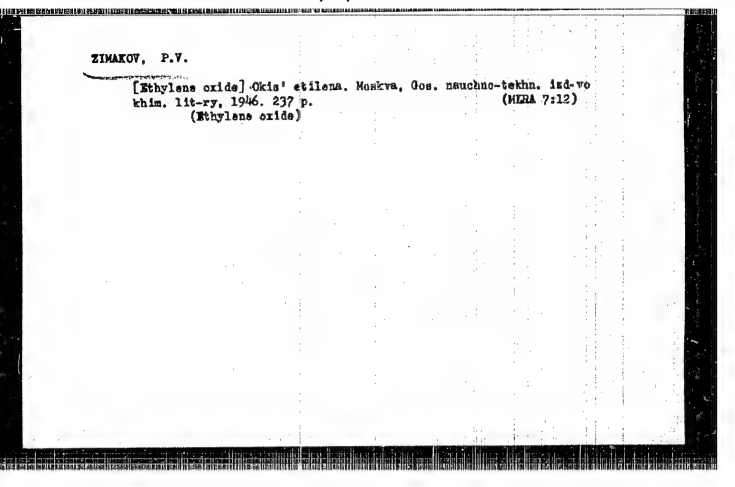


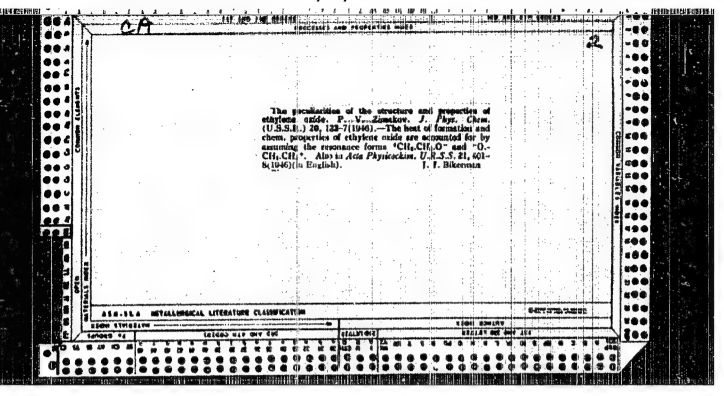


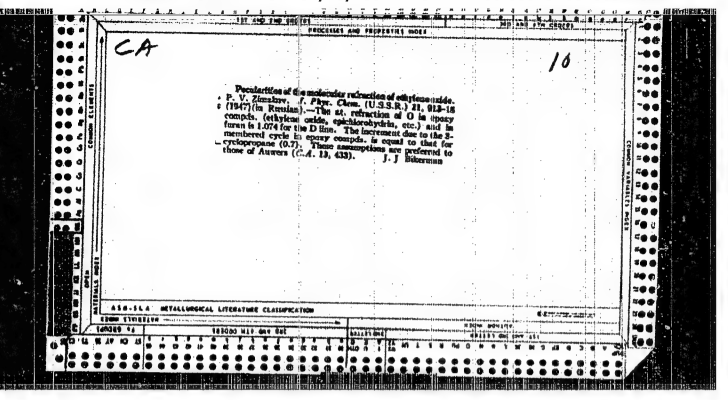


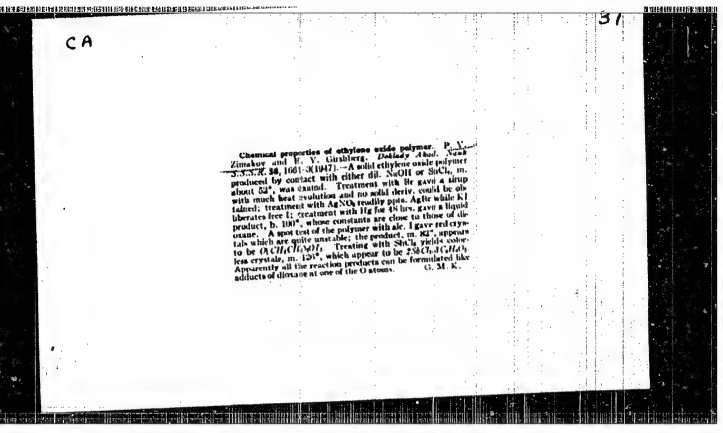


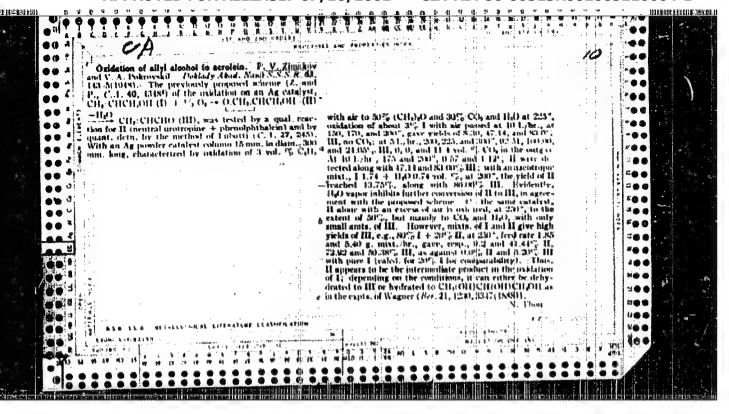


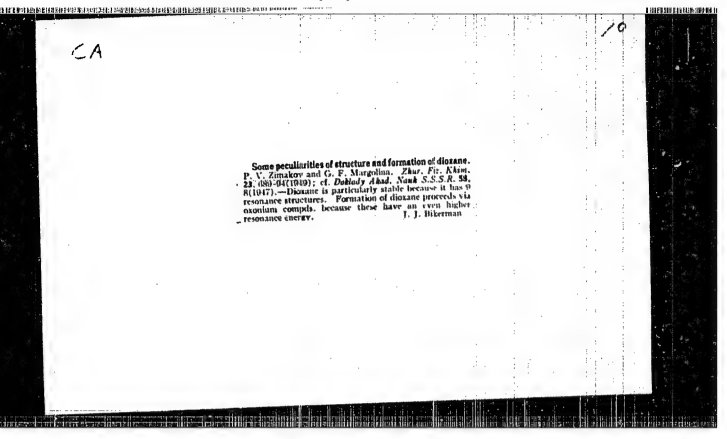


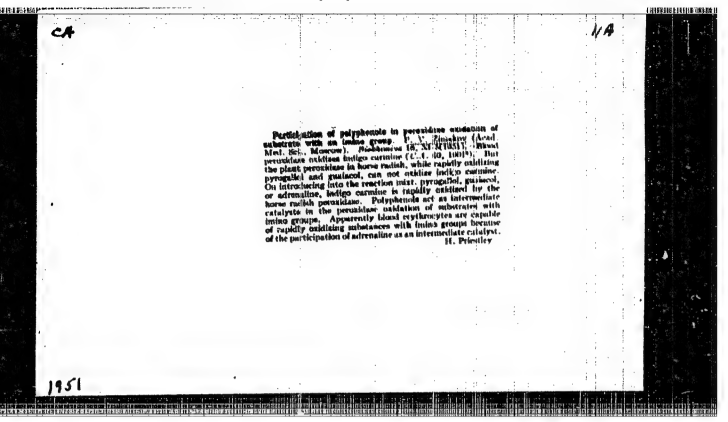


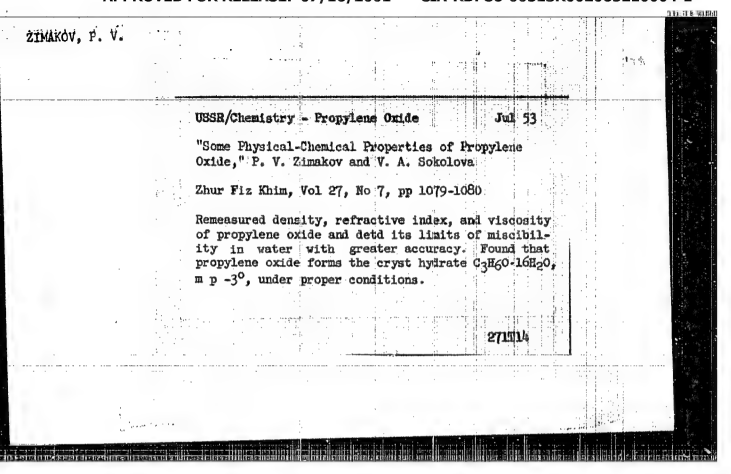


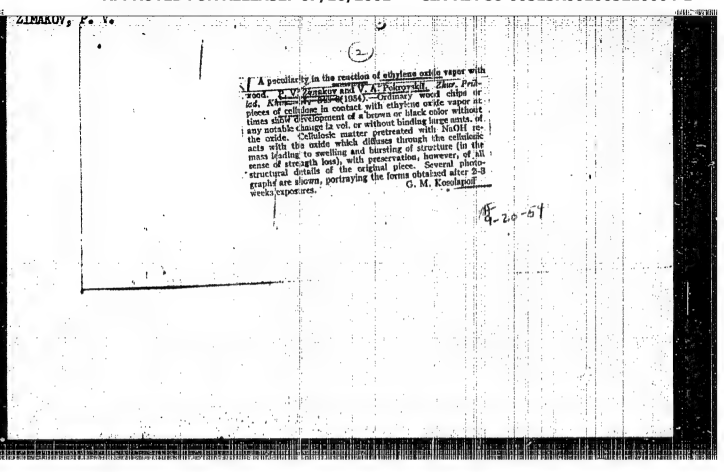


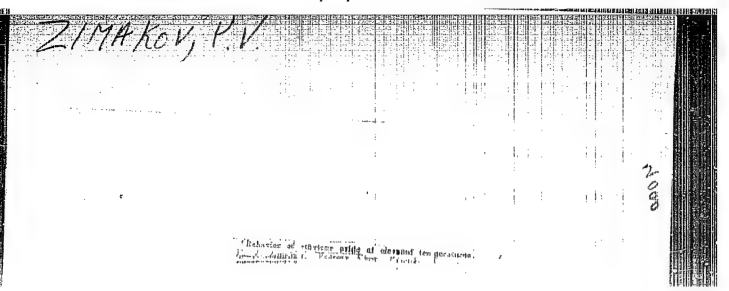


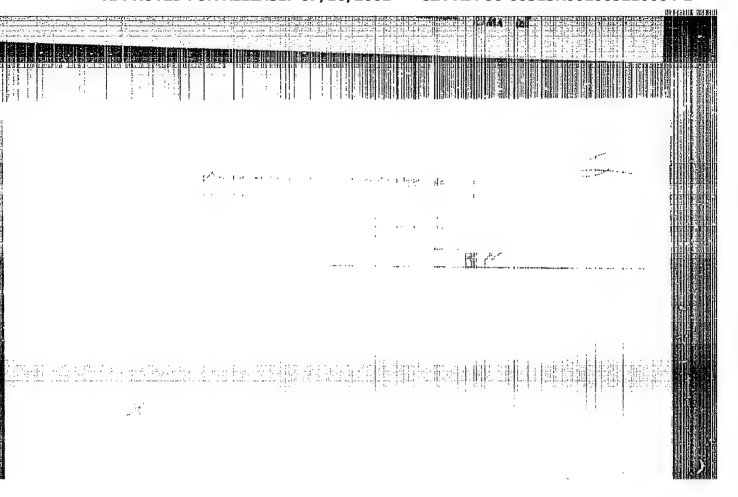




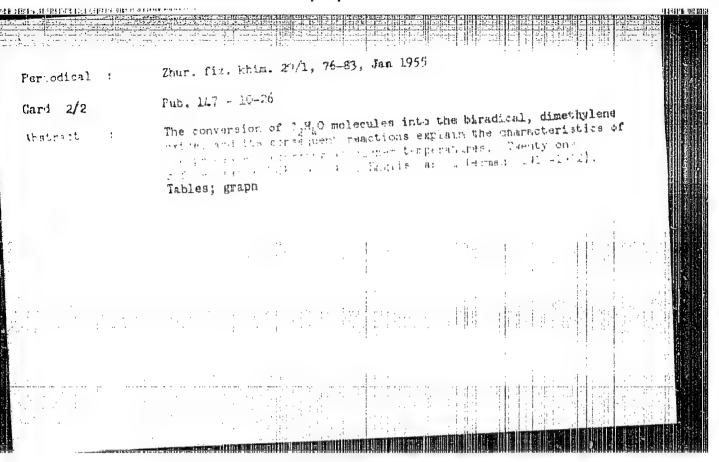






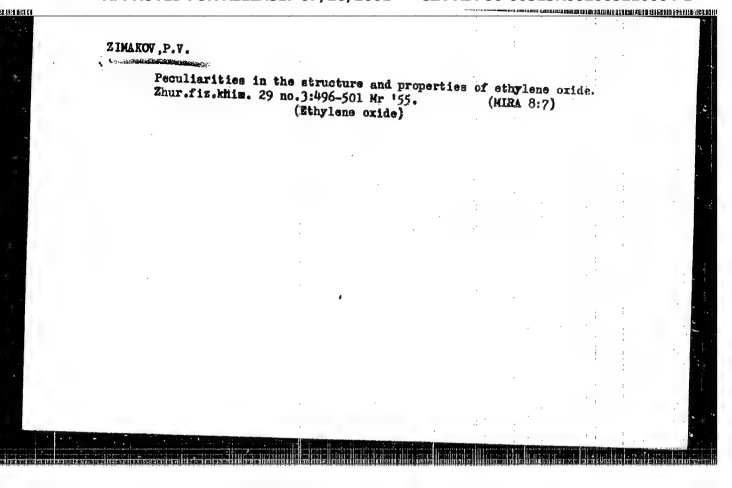


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CIA-RDP86-00513R002065210004-1



ZIMAKOV, P.V.

Category: USSR / Physical Chemistry-Molecule. Chemical bond

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Abs Jour: Referat Zhur-Khimiya, No 9, 1957, 29552

Author : Zimakov, P.V. Inst : not given

Title : Experimental Confirmation of the Presence of Oxonium Form of

Ethylene Oxide in Aqueous Solutions

Orig Pub: Zh. fiz. khimii, 1956, 30, No 8, 1904-1905

Abstract: Distortion of valency angles in the molecule of ethylene oxide (I) must increase the affinity of oxygen to the proton and its tendency to conversion to the "oxonium" state. Comparison of blowing with air, at a rate of about 10 liters per hour, of 1.5-1.6% solutions of I, in neutral and in acidified water, at 15-20°, has revealed that from neutral solutions I is driven of completely within 15 minutes, while in acidified solutions up to 15% of the initial amount of I are always retained. In the opinion of the author this

is due to a lowering of the vapor tension of I, caused by a solvation, catalyzed by H ions, with formation of the exemium form of I.

Card : 1/1

-5

ZIMAKOV, USSR/Chemical Technology - Chemical Products and Their Application. Industrial Organic Synthesis. : Ref Zhur - Khimiya, No 1, 1958, 2146 Abs Jour : Zimakov, P.V. Author Inst Current Methods for the Preparation of Ethylene Oxide Title : Khim. nauka i prom-st', 1957, 2, No 1, 24-33 Orig Pub : A review article. Considered in detail are the two princi-Abstract pal current methods for the preparation of ethylene oxide (I), which are of significance in the industrial production of I: a) the method comprising the step of hypochlorination of ethylene (II) with subsequent decomposition of the result mg HO(CH2);Cl, and b) the method of direct oxidation of II, with 0, or air, to I, without catalyst (volumetric oxidation) or over an Ag catalyst. Views concerning the mechanism of both processes are considered and discussed. A comparison is made of the advantages and Card 1/2

ZIMAKOV, P.V.

20-2-28/62

AUTHOR TITLE ZIMAKOV, P.V., and KOGAN, L.M.

On Two Reaction Trends in the Dehydrochlorination of Lower Alkylene

Chlorohydrins

(O dvukh napravleniyakh reaktsii degidrokhlorinovaniya mizshikh alkilen-

khlorgidrinov, Russian)

Doklady Akademii Nauk SSSR, 1957, Vol 115, Nr 2, pp 297-300 (U.S.S.R.)

PER IODICAL ABSTRACT

In the year 1959 it will be hundred years since the description by A. Wirz of the reaction mentioned in the title. At that time a new compound, ethylene chlorids, was for the first time produced in an alkaline medium. At present this reaction forms the basis of the commercial production of ethylene by means of chlorine. No note was hitherto taken of the fact that this substance is always obtained with an admixture of $1-2^{\circ}/_{\circ}$ acetaldehyde, if the reaction is performed with application of milk of lime. These admixtures can develop due to a secondary isomerization reaction of ethylene oxide. In 1939 a French patent was published concerning the formation of considerable quantities of alderhydes (or ketones) beside the gravides on dehydrochlorination of alimpathic chlorohydrins in the environment of "milk of Magnesia". The subject of the patent did not draw the attention of chemists to the two-way reactivity of the simple compound (ethylene chlorohydrin). Some years ago the authors made thorough investigation of the dehydrochlo-

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and the state of t

On Two Reaction Trends in the Dehydrochlorination of Lower Alkylene Chlorohydrins

rination of ethylenechlorohydrin (denoted ECIH in the following) in various media and conditions. The method of the French patent yielded up to 35 % of ethyleneoxide and up to 50 % acetaldehyde. The authors proved that ethylene glycol develops only in the hydratation of the primarily-forming ethyleneoxide. Acetaldehyde, as the practically only reaction product $(98^{\circ}/_{\circ})$, with small admixtures of ethyleneoxide develops in a boiling water suspension of nickeloxide hydrate. This proves that even such a simple compound as ECIH, according to the nature of reagents acting on it, distinctly shows two reaction trends. It was interesting to study the particularities of the dehydrochlorination of ECIH. In order to make the experimental checking of the above-mentioned assumption (isomerization) easier, the authors used propylenechlorohydrin (=PClH), since this compound exists in 2 (α- and β-) isomeric forms. Its dehydrochlorination takes place just as easily as that of ECIH, but it usually leads to a yield of propylene oxide higher than 85%. The causes of the small yield have not been determined. Conclusions: 1.) Propion-aldehyde in greater quantities develops on dehydrochlorination of PClH only in suitable media: aqueous suspensions of magnesium- and especially nickel-hydrooxide. This is in agreement with the rule governing in the case of ECIH. 2.) Propion aldehyde de-

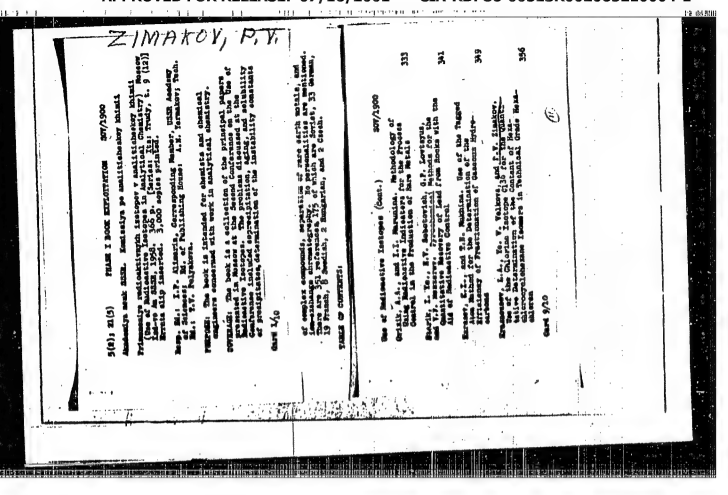
Card 2/3

ZIMAKOV, P. V., BYKOV, A. G. and USACHEVA, I. A. (Ministry of the Chemical Industry)

"Radio Electrochromatographic Method of Analysis"

Isotopes and Radiation in Chamistry, Collection of papers of 2nd All-Union Sci. Tech. Conf. on Use of Radioactive and Stable Isotopes and Radiation in Mational Economy and Science, Moscov, Ezd-vo AM SSUR, 1958, 180pp.

This volume published the reports of the Chemistry Section of the 2nd AU Sci Tech Conf on Use of Radioactive and Stable Isotopes and Radiation in Science and the National Recommy, apondored by Amad Sci. USER and Main Admin for Utilization of Atomic Energy under Council of Ministers USER Moscow 4-12 Apr 1957.



ZIMAKOV, P. V. and KRASNOUSOV, L. A.

"Use of C1³⁶."

report presented at The Use of Radioactive Isotopes in Analytical Chemistry, Conference in Moscow, 2-4 Dec 1957

<u>Yestnik Ak Nsuk SSSR</u>, 1958, No. 2, (author Rodin, S. S.)

507/ 64-58-4-5/20

AUTHORS:

Zimakov, P. V., Doctor of Chemical Sciences, Kogan, L. M.,

Candidate of Technical Sciences

TITLE:

On the Influence of Temperature on the Process of the Hypochlorination of Ethylene (O vliyanii temperatury ma

proteess gipokhlorirovaniya etilena)

PERIODICAL:

Khimicheskaya promyshlennost', 1958, Nr. 4, pp. 210 - 213(USSR)

ABSTRACT:

Already Carius (Ref 1) observed the reaction between ethylene and hypochlorous acid, however, a technological use of this reaction took place only according to the synthesis by Gomberg (Ref 2). Greatly differing data exist on the influence of the temperature on the course of reaction: Brooks (Ref 21) maintained that the hypochlorination is to be carried out at low temperatures; Zapadinskiy (Ref 3) worked at 0 .. 2 while Tropsch and Kassler (Ref 10) showed that the reaction takes a better course at 40 - 60 than at 0 - 30. According to Domask and Kobe (Ref 15) a rise of temperature is to supply a better yield, while Murray (Ref 24) finds 50 to be the upper limit. These contradictions made the authors

Card 1/4

SOV / 64-58-4-5/20 Hypochlorination

On the Influence of Temperature on the Process of the Hypochlorination of Ethylene

of this paper assume that the distribution of ethylene and with it its transformation velocity into the solution are dependent on temperature. For this purpose three experimental series were carried out at temperatures of from 0 - 70 and in the case of different acetylene distribution; in the first case a distribution of the gas was arranged by a Schott filter Nr 1, in the second case by openings in the supply tube, and in the third case by means of 1 mm openings. From the results obtained may be seen that in the first series of experiments a rise of temperature lead; to an increase in the yield of ethyleme chiorohydrin (at 70 up to 90 %). The second series showed that until three conditions a change of temperature within a wide range did not exert any influence on the yield and that on the average it amounts to 20 %. In the third series of experiments a rise of temperature caused a decrease of the ethylene chlorohydrin yield, so that it was 40 % at 90° and 55 % at 60°. In order to explain these contradicting results the process of hypochlorination is shown schematically and two basic reactions are assumed - the solution of ethylene and the chlorohydrolysis - which influence the yield. The hydrolysis was already investigated

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On the Influence of Temperature on the Process of the Hypochlorination of Ethylene

by Yakovkin (Ref 25) who observed an increase in the degree of hydrolysis with the rise of temperature; for the further classification an investigation of the solution kinetics is carried out. It is assumed that for the first experimental series a "kinetic" factor is decisive, while the third experimental results depend on a "diffuse" factor, and the second series has a balancing effect of the temperature on the two mentioned factors; such an independence of the temperature may be observed after all. Based on the results obtained the authors then conclude that a rise of temperature has a favorable effect in the case of a fine distribution of the gases, while in a coarse distribution the temperature factor exerts a negative effect. Thus for obtaining a maximum yield of ethylene chlorohydrin the corresponding conditions must be prepared; the effect of the optimum temperature will the velocity of the transition be the greater, the greater of ethylene from the gaseous phase into the solution - the medium of the meaction. There are 1 figure and 25 references, to of which are Soviet.

Card 3/4

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KRASHOUSOV, L.A.; VOLKOVA, Ye.V.; ZIMAKOV, P.V.

Chlorine isotope Cl³⁶ used for quantitative determination of the isomeric composition of hexachlorocyclohexane in commercial hexachloran. Trudy kom.anal.khim. 9:356-360 158. (MIRA 11:11) (Cyclohexane) (Radioactive tracers) (Chlorine—Isotopes)

ZIMAKOV, P.V.

"Some questions of the fixation of radioactive isotopes in connection with the problem of their safe burial."

report presented at the Scidntific Conference on the Disposal of Radioactive Wastes, Monaco, November 1959.

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5 (3) AUTHORS:

Zimakov. P. V. Kogan, L. M.

SOV/20-127-2-26/70.

TITLE:

The Mechanism of Aldehyde Formation in the Dehydrochlorination of Lower Alkylene Chlorohydrins

PERIODICAL:

Doklady Akademii nauk SSSR, 1959, Vol 127, Nr 2, pp 329-332 (USSR)

ABSTRACT:

The authors proved (Ref 1) that the transformation of the substances last mentioned in the title (sthylene-, propylene chlorohydrin) into a corresponding co-oxide or aldehyde with good yields depends on the character of the medium in which the reaction proceeds. However, only one isomer, i.e. the \$-isomer, of propylene chlorohydrin is transformed into an aldehyde, the cisomer is not capable of being transformed (see Scheme). The transformation of the mentioned substances into c-oxides proceeds in media with a high pH according to the Wirth-(Vyurts-) reaction (in "milk of lime"). They are transformed especially smoothly into aldehydes in a suspension of nickel hydroxide (Ref 1). The aldehyde yields in "milk of magnesia" are as well considerable (Ref 2). The authors give then considerations with respect to the probable mechanism of such a double-track reactivity of such simple compounds as those mentioned in the title. They have two easily polarisable Cl- and O-atoms which are in a

Card 1/4

The Mechanism of Aldehyde Formation in the Dehydrochlorination of Lower Alkylene Chlorohydrins

BOY/20-127-2-26/70

β-position to each other. This favors a maximum reciprocal effect which is often accompanied by the change of the valence stages of both atoms or of one of them. The yperite melecule may serve as an example of such an "activity" of the mentioned &-pairies of the chlorine atoms and of an easily polarisable sulphur. Its very special toxicity is explained by the slight change in the valence of the mentioned atoms (Ref 3). The activity of the polarizable atoms is also very high in the two mentioned chlorohydrine in consequence of their structural peculiarities. The reactivity is due to this fact. Even an anhydrous ethylene chlorohydrin produces a certain quantity of diethylane-glycolchlorohydrin in the case of a long storage. An equivalent HOL-quantity becomes liberated in this case (Ref 4). This transformation is caused by the transition of single chlorohydrin molecules into an "active" polar form with changed valence stages of oxygen and chlorine (Ref 5). The Wirts reaction which leads to the formation of ethylene-oxide proceeds through such an active molecule form of chlorohydrin with exemium exygen and with ion chlorine (Ref 6). The propylene cride is produced from propylene ohlorohydrin in the same way. Its two isomers (at and 6)

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The Mechanism of Aldehyde Formation in the Dehydrochlorination of Lower Alkylene Chlorohydrins

react practically equally. The ethylene-oxide produced in the practical carrying-cut of the Wirtz reaction is: a 1 w a y s accompanied by a small admixture of acetaldehyde (Ref 7). The accompanied by a small admixture of acetaldehyde (Ref 7). The formation of the latter increases according to reference 8 with decreasing acidity of the medium. At present it must be assumed that the ethylene chlorohydrin molecule may assume a second active "chloronium" form under splitting off of a hydroxyl ion in media "chloronium" form under splitting off of a hydroxyl ion in media not containing an excess of hydroxyls. This must be assumed from the ethylene chlorohydrin formation in the ethylene chlorination in the aqueous medium (Ref 10). Analogous active forms exist for propylene chlorohydrin, however, with a different degree of stability (Ref 11). Nickelous hydroxide N(OH) in suspension is

an especially suitable medium, as was already mentioned (Ref 1). It may be assumed that the interaction of ethylene chlorohydrin in the aqueous medium passes in the case of boiling (100°) several stages explained in the scheme. Such alcoholate forms of the nickel compounds are described in reference 13. The considerable tendency of nickel to the formation of a basic chloride is essential, whereas the intermediate compound with

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The Mechanism of Aldehyde Formation in the Dehydrochlorination of Lower Alkylene Chlorohydrine SOV/20-127-2-25/70 alkylene chlorohydrin favors the reconstruction of the molecule of the latter under splitting off of an acetaldehyde. The «-isomer of propylene chlorohydrin which does not incline to the splitting off of hydroxyl and to the formation of chloronium does not react at all with the nickelous hydroxide (Ref 1). The two chains of the interactions are simultaneous and parallel in the "milk of magnesia" in which the yield of Oc-oxides and aldehydes is approximately equal (Ref 14). There are 14 references, 8 of which are Soviet. PRESENTED: April 1, 1959, by B. A. Kazanskiy, Academician SUBMITTED: March 30, 1959 Card 4/4

ZIMAKOV, P. V., ZAKHAROVA, K. P., KULIHHENKO, V. V., HOGDANOV, N. I. (USSR)

"A Thermic Method of Preparing Sr-90 Sources."

report presented at the Conference on Radioisotopes in Metallurgy and Solid State Physics, IAEA, Copenhagen, 6-17 Sept 1960.

s/089/60/009/005/012/020 B006/B070

AUTHORS:

Krasnousov, L. A., Zimakov, P. V., Volkova, Ye. V.

TITLE:

Radiochemical Chlorination of Benzene

PERIODICAL:

Atomnaya energiya, 1960, Vol. 9, No. 5, pp. 412 - 414

TEXT: The radiative chlorination of benzene was studied under standard conditions in order to study the possibility of using nuclear radiations for the production of hexachlorane. As can be seen from the Table, the different radiations led to the formation of hexachloracyclohexane (HCCH) characterized by a high content of alpha phase. In addition to data on thermal, chemical (benzene peroxide), and infrared chlorination, the Table gives the following data:

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Concentration g chlorine/100g C6H6	t°C	γ-Isomer %	a-Isomer %	Mean energy ev
Ultraviolet (3650A)14.0 β: (3.48 rad/sec,	40	11.3	.73.8	3 - 5
total: 2.4.10 ⁴ rad)14.0 y: (6.75 rad/sec,	40	10.2	78.5	0.4.106
total: 2.4.10 ⁴ rad)14.0	40	11.8	83.5 1	2-106-1-3-106
On the contrary, the content only 63.7%. The chlorination tion in CCl ₄ . The β source	n was d	lone for pur	e substance	and for solu-
chemical yield of the react strongly dependent on the p ally pure benzene is used, yielded 9000 molecules per	urity of the yie 100 ev.	of the start old is only . The radiat	ing material 130,000. The ive chloring	l. If industri- e ultraviolet ation rate is

proportional to the square of radiation intensity (benzene without solvent). In CCl solution, the rate of reaction is essentially lower.

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Radiochemical Chlorination of Benzene

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Radiative chlorination of benzene is possible also at low temperatures in solid phase; the total yield increases with decreasing temperature down to -80° C. Chlorination remains incomplete for temperatures still lower (at -120° C, the content of tetrachlorocyclohexane reaches 58%). The effect of temperature on the isomeric composition of HCCH was also studied. While the total yield of HCCH has its maximum at -80° C, the content of α -isomer decreases from 83.5 to 38% for the fall of temperature from 40 to -190° C. The yield of γ -isomer also depends on the concentration of chlorine. The formation probabilities of α -, β -, γ -, δ -, and \mathcal{E} -isomers were calculated to be 27.8, 4.63, 25.0, 26.0, and 16.7%, respectively. There are 4 figures, 1 table, and 3 references: 1 Soviet, 1 German, and 1 Polish.

SUBMITTED:

March 31, 1960

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35/37 5/081/62/000/004/074/087 B138/B110

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AUTHORS:

Zimakov, P. V., Volkova, Ye. V., Fokin, A. V., Sorokin, A. D

Belikov, V. M.

TITLE: Use of nuclear radiation energy in the process of the

polymerization of fluoro-olefines

PERIODICAL: Referativnyy zhurnal. Khimiya, no. 4, 1962, 557, abstract 4P24 (Sb. "Radioakt. izotopy i yadern. izlucheniya v nar.

kh-ve SSSR, v. 1. M.", Gostoptekhizdat, 1961, 219-226)

TEXT: The processes of the separate and combined radiation polymerization of tetrafluorethylene and trifluorchlorethylene have been investigated with the aim of eliminating some of the deficiencies in existing methods of fluoro-olefine polymerization. It has been found that tetrafluorethylene and trifluorchlorethylene can easily be polymerized under various temperature conditions and mediums with comparatively low radiation intensities. The resulting polymers have a high degree of purity. The possibility of producing various fluoro-copolymers by radiation is demonstrated. Both radiation polymerization and radiation vulcanization might be carried out in the case of fluor-containing rubbers. [Abstracter's note: Complete translation.]

8/089/64/010/001/008/020 B006/B063

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Zimakov, P. V., Kulichenko, V. V.

TITLE:

Some Problems Concerning the Localization of Radioisotopes

in Connection With the Problem of Their Safe Storage

Atomnaya energiya, 1960, Vol. 10, No. 1, pp. 58-63 PERIODICAL:

TEXT: The authors have made a detailed study of the methods used today for the storage of radioactive waste matter, and they now discuss the hazards involved. It must be borne in mind that, in general, radioactive wastes will actually remain on the spot for several centuries, especially if it contains Sr90, Cs137, and similar isotopes. First, the authors reject the widespread opinion that fluid radioactive wastes can be safely stored in any container. Apart from corrosion, there may arise considerable overpressure in the gas container. This overpressure results from radiolytically evolving gases and might lead to the destruction of the container. Certain radioactive solutions are capable of evolving gas in quantities of up to 10 cm3/cure per hour. In addition, the activity of the waste matter may heat the container and thus destroy it through evolution of

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Some Problems Concerning the Localization of Radioisotopes in Connection With the Problem of Their Safe Storage

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vapor or pyrochemical processes. Storage at great depths does not prevent radioactive fluids from penetrating into the ground water. The most promising method is to solidify all fluid radicactive matter, a possibility that is discussed in detail. Of special interest is the conversion of radioactive wastes into difficultly soluble precipitates, such as hydroxides, phosphates, etc. The best way to keep radioactive wastes on the spot is to deposit isotopes in the form of vitreous preparations of the smallest possible size. An analysis of the physicochemical fundamentals of producing such preparations is presented, and some specific features of the state and behavior of sealed-in radioactive fission fragments are discussed. The melting processes and also the formation of radioactive aerosols (which increases rapidly with temperature, especially above 1200°C) in the heat treatment of radioactive slimes are described. Fractional and X-ray structural analyses have shown that the melts obtained are inhomogeneous, i.e., the vitreous, amorphous preparation contains crystalline inclusions, particularly iron compounds, which are the principal carriers of radioactive fragments. Problems of leaching out and elution of radioactive matter by ground water, as well as self-heating

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